Oncology Clinical Pathways Breast Cancer

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Breast Cancer – Presumptive Conditions

VA automatically presumes that certain disabilities were caused by military service. This is because of the unique circumstances of a specific Veteran's military service. If a presumed condition is diagnosed in a Veteran within a certain group, they can be awarded disability compensation.

Atomic Veterans Exposed to Ionizing Radiation

Breast cancer

Gulf War and Post 9/11 Veterans

If the patient served on or after Sept. 11, 2001, in Afghanistan, Djibouti, Egypt, Jordan, Lebanon, Syria, Uzbekistan, or Yemen or if the patient served in the *Southwest Asia theater of operations, or Somalia, on or after Aug. 2, 1990, specific conditions include:

Reproductive cancers of any type

* The Southwest Asia theater of operations refers to Iraq, Kuwait, Saudi Arabia, the neutral zone between Iraq and Saudi Arabia, Bahrain, Qatar, the United Arab Emirates, Oman, the Gulf of Aden, the Gulf of Oman, the Persian Gulf, the Arabian Sea, the Red Sea, and the airspace above these locations.

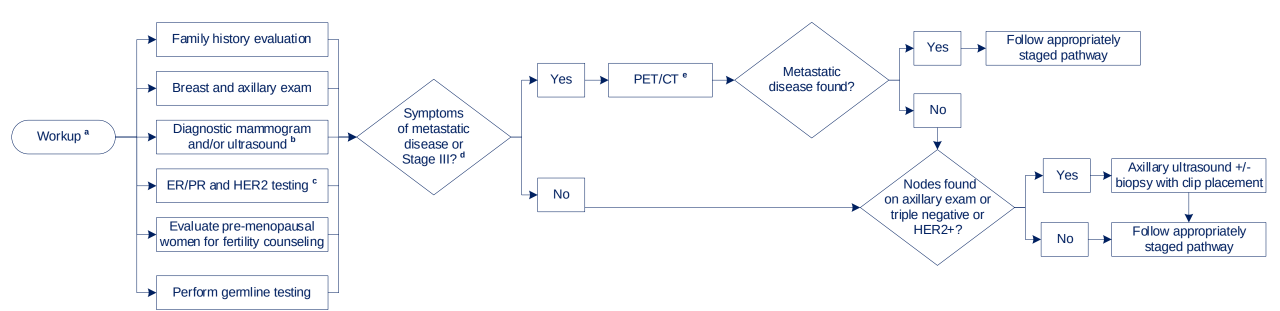
For more information, please visit U.S. Department of Veterans Affairs - Presumptive Disability Benefits (va.gov)







Breast Cancer – Workup



Clinical trial(s) always considered on pathway.

- ^a Workup after biopsy-proven invasive cancer
- Diagnostic Imaging if not previously performed; MRI not routinely recommended
- ER/PR and HER2Testing follow Pathology pathway for in-depth information
- d Metastatic Disease confirmation by biopsy; symptoms include neurological symptoms, persistent cough, abnormal blood counts, abnormal LFTs, bone pain; if neurological symptoms, perform brain MRI with contrast
- e PET/CT if unavailable, perform CT chest/abdomen/pelvis with bone scan







Breast Cancer – DCIS



Clinical trial(s) always considered on pathway.

^a ER testing is recommended; HER2 testing is not recommended

b Breast Conservation ineligibility includes inability to obtain clear margins without mastectomy or patient is not candidate for radiation

conservation?

^c Lumpectomy sentinel node biopsy may be recommended based on high grade, palpable tumor, anatomic location compromising future sentinel lymph node, or extensive volume

Plastic Surgery

^d ER Positive if staining ≥ 1% by IHC

ER Negative if staining < 1% by IHC</p>

Menopausal defined as patient that is \geq 60 years of age; \geq 1 year amenor thea (not medically induced); history of Bilateral Salpingo-Oophorectomy (BSO); or confirmed with labs

⁹ Tamoxifen avoid tamoxifen if prior history of DVT or known hypercoagulability

h Anastrozole evaluate baseline bone density; promote weight-bearing exercise, smoking cessation, reduced alcohol intake, and calcium/vitamin D supplementation





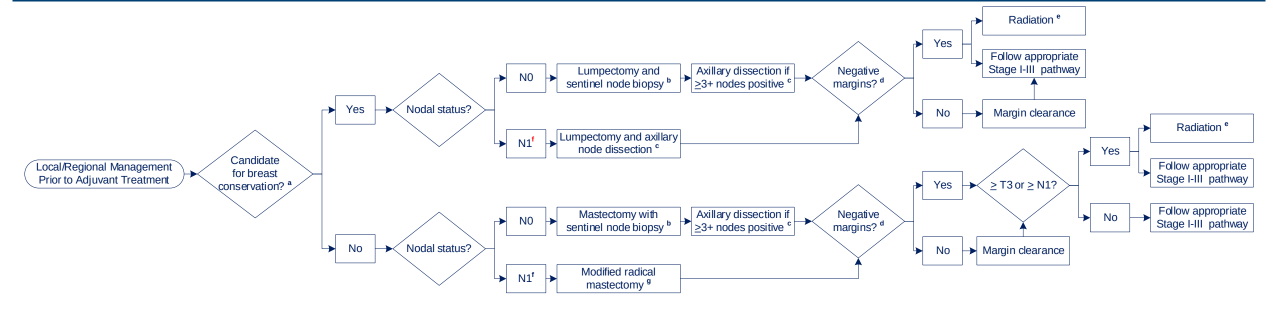
Unilateral mastectomy and Refer to Medical Oncology

for risk reduction discussion

sentinel node biopsy



<u>Breast Cancer – Local/Regional Management Prior to Adjuvant Treatment</u>



Clinical trial(s) always considered on pathway

^a Breast Conservation ineligibility includes inability to obtain clear margins without mastectomy, or patient is not candidate for radiation; if mastectomy early referral to Plastic Surgery is recommended; if lumpectomy early referral to Radiation Oncology is recommended; same treatment for male patients, however it is recognized that the majority of male patients will elect for mastectomy

^b Sentinel Node Biopsy not routinely recommended if patient age > 69 and T1 ER+/HER2- tumors

Axillary Dissection includes complete level I/II clearance

d Negative Margins defined as no tumor on ink

e Radiation if patient <T2 and <2 positive nodes patient can opt for nodal radiation in lieu of axillary dissection; in patients where (only) whole breast RT is planned, hypofractionated treatment is preferred over conventional fractionation; in select cases Accelerated Partial Breast Irradiation (APBI) is an acceptable treatment option

fN1 Disease recommend neoadjuvant chemotherapy include HER2+ and TNBC patients

⁹ MRM includes axillary dissection

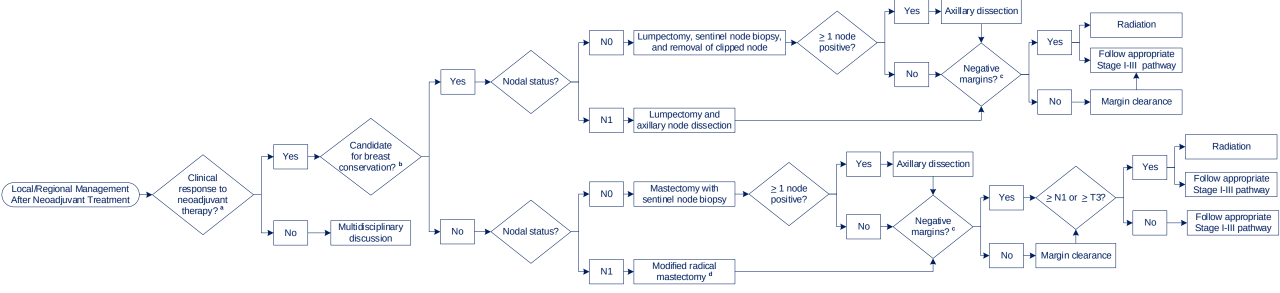
MRM Modified Radical Mastectomy
TNBC Triple Negative Breast Cancer







<u>Breast Cancer – Local/Regional Management After Neoadjuvant Treatment</u>



Clinical trial(s) always considered on pathway

a Clinical Response determined by exam and/or imaging

Preast Conservation ineligibility includes inability to obtain clear margins without mastectomy or patient is not candidate for radiation; early referral to Plastic Surgery is recommended

Negative Margins defined as no tumor on ink

MRM includes axillary dissection

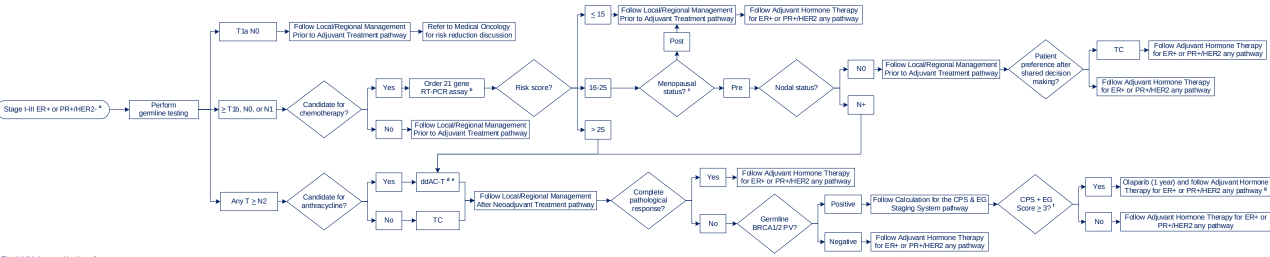
MRM Modified Radical Mastectomy







Breast Cancer – Stage I-III ER+ or PR+/HER2-



Clinical trial(s) always considered on pathway.

^a Invasive Carcinoma to include ductal, lobular, metaplastic, and mammary; less aggressive breast carcinoma, actionoma, mucinous (colloid) carcinoma, mucinous cystadenocarcinoma, acerciony carcinoma, acerciony carcinoma, acerciony carcinoma, and tall cell carcinoma with reversed polarity

b Blocks Preferred to Unstained Slides if using unstained slides, one must submit 15 5-um-thick sections that are numbered to indicate their order, choose tissue from the block with the greatest contiguous area of the highest grade of invasive carcinomas are not acceptable; biopsy, lumpectomy, and resection specimens can be used; tissue must have been fixed in formalin

Menopausal defined as patient that is > 60 years of age. > 1 year amenorrhea (not medically induced), history of Bilateral Salpingo-Oophorectomy (BSO), or confirmed with labs

d ddAC-T followed by weekly paclitaxel (T)

e Evaluate Cardiovascular Risk factors with baseline LVEF and CMP

CPS + EG Score incorporates estrogen receptor (ER) status and tumor grade with pretreatment clinical stage (CS) and post-treatment pathologic stage (PS); Follow Calculation for CPS & EG Staging System pathway for further information

⁹ Olaparib patients should not be on concomitant olaparib and abemaciclib therapy

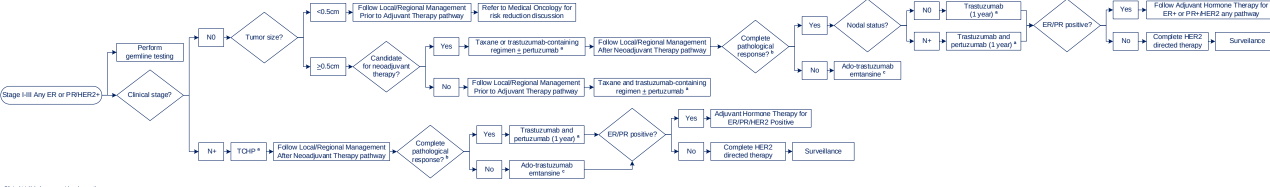
CMP Comprehensive Metabolic Panel
ddAC-T Dose-dense AC-T (doxorubicin and cyclophosphamide)
LVEF Left Ventricular Ejection Fraction
PV Pathogenic Variant
TC docetaxel and cyclophosphamide







Breast Cancer – Stage I-III Any ER or PR/HER2+



Clinical trial(s) always considered on pathway

a Evaluate Cardiovascular Risk Factors with baseline LVEF (with ECHO or MUGA) and CMP; monitor LVEF every 3 months during therapy

^b Complete Pathological Response absence of residual invasive carcinoma in both the breast and lymph nodes

^c Ado-trastuzumab Emtansine radiation and hormone therapy can be given concomitantly with trastuzumab, pertuzumab, and ado-trastuzumab emtansine

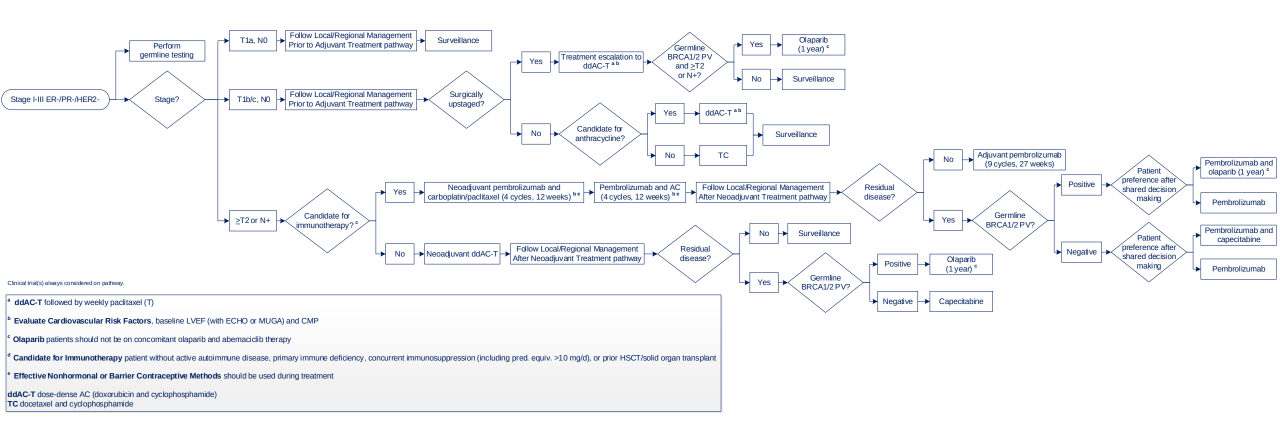
TCHP docetaxel/carboplatin/trastuzumab/pertuzumab







Breast Cancer – Stage I-III ER-/PR-/HER2-

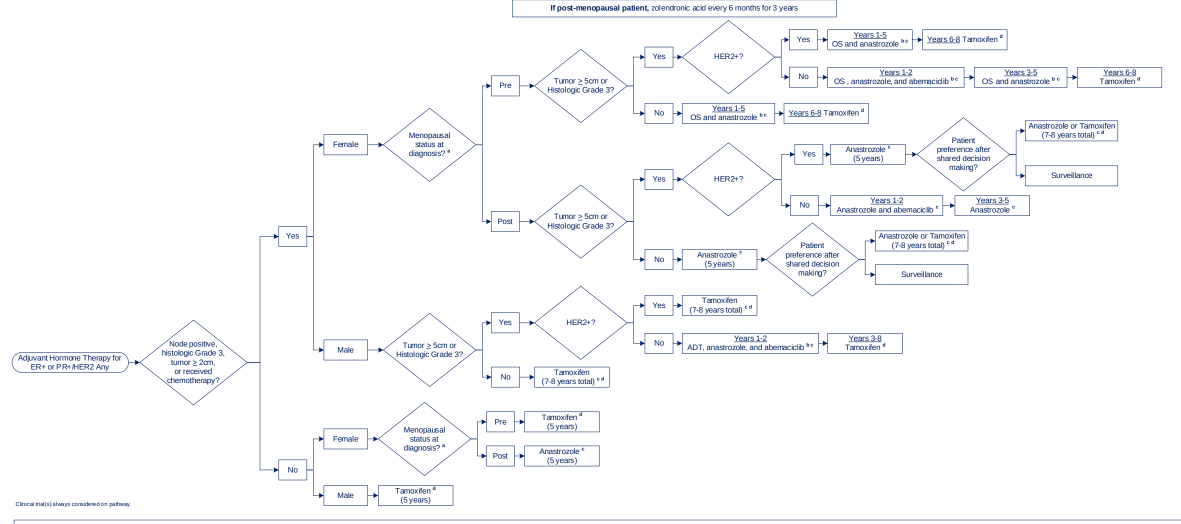








Breast Cancer – Adjuvant Hormone Therapy for ER+ or PR+/HER2 Any



a Menopausal defined as patient that is ≥ 60 years of age; ≥ 1 year amenorrhea (not medically induced); history of Bilateral Salpingo-Oophorectomy (BSO); or confirmed with labs

Ovarian Suppression (OS) includes surgical or medical suppression

⁶ Anastrozole only for post menopausal women or women undergoing ovarian suppression; evaluate baseline bone density; promote weight-bearing exercise, smoking cessation, reduced alcohol intake, and calcium/vitamin D supplementation; if not a candidate for anastrozole, tamoxifen is an alternative; if patients do not tolerate one AI, any AI is a suitable alternative

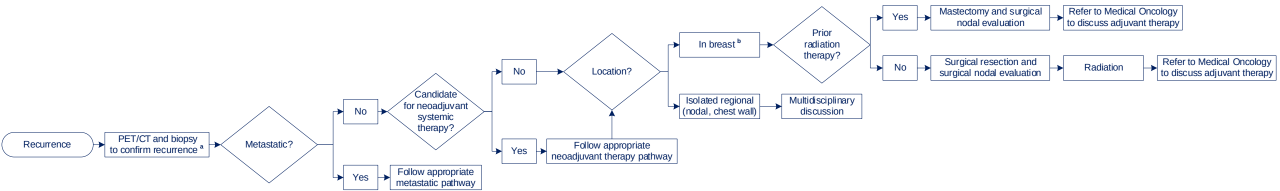
^d Tarnoxifen avoid tamoxifen if prior history of DVT or known hypercoagulability; if contraindication to tamoxifen in men, prescribe AI with ADT; patients should use effective nonhormonal contraception or barrier contraceptive during tamoxifen therapy; continue for 2 months after last dose







Breast Cancer – Recurrence



Clinical trial(s) always considered on pathway.

^a **PET/CT** if unavailable, perform CT chest/abdomen/pelvis with bone scan

^b Multidisciplinary Discussion highly recommended for this patient presentation







Breast Cancer – Stage IV ER+ or PR+/HER2-

If patient is in visceral crisis (imminent organ failure), proceed to chemotherapy; if disease becomes stable, resume endocrine therapy Fulvestrant and alnelising and OS or ADT d Perform germline testing and CGP testing AKT, PTEN Anastrozole and ribociclib and OS or ADT de ≥12 months Germline BRCA 1/2 PV, somatic BRCA Olaparib since prior adjuvan Molecular target Stage IV ER+ or PR+/HER2- a b c 1/2 PV, or germline PV in PAI B2 therapy or newly identified? diagnosed? Trastuzumab deruxtecan g h and ribociclib of Everolimus and exemestane and OS or ADT ° HER2 low Prior fulvestrant? ESR1? Everolimus and fulvestrant line of prior and OS or ADT ° chemotherapy Candidate for Received No ≥ 2 prior lines of chemotherapy? Paditaxel ^j Hormonal-based therapy ^a If Bone Metastases prescribe zoledronic acid; prescribe denosumab if contraindication; vitamin D and dental evaluation; if symptomatic, refer to Radiation Oncology Capecitabine 1 ^b If Brain Metastases referral to radiation oncology; refer to Neurosurgery olf MSI High or TMB ≥10 pembrolizumab is recommended to be given after endocrine therapy Carboplatin 1 d Ribociclib risk of QTc interval prolongation: evaluate ECG at baseline, regular intervals and as clinically indicated, ensure electrolyte abnormalities are corrected prior to start; monitor for neutropenia, hepatobiliary toxicity, cutaneous adverse reactions; reduce initial dose if renal and/or hepatic impairment Eribulin " OS and/or ADT ovarian Suppression (OS) must be added for women who are pre-menopausal or androgen deprivation therapy (ADT) for men Capivasertib evaluate fasting blood glucose and HbA1C prior to start and at regular intervals; closely monitor for diarrhea and cutaneous adverse reactions; note odd dosing: 4-days on then 3-days off Gemcitabine ⁹ Evaluate Cardiovascular Risk Factors with baseline LVEF (with ECHO or MUGA) and CMP; monitor LVEF every 3 months during therapy Trastuzumab Deruxtecan has proven overall survival advantage; one prior chemotherapy in the metastatic setting; monitor for interstitial lung disease Sacituzumab Govitecan has proven overall survival advantage; after two chemotherapies or one chemotherapy in the metastatic setting if adjuvant chemotherapy was given within the past 12 months; monitor for diarrhea and cytopenias Ixabenilone Vinorelhine 9 Capecitabine if oral agent is preferred; less hair loss as compared to other agents Carboplatin if not used in 1L setting; preferred in women with BRCA pathogenic variants Pembrolizumab m Eribulin monitor for neuropathy and cytopenias Liposomal Doxorubicin baseline LVEF > 50% and/or no clinically significant cardiac disease Vinorelbine monitor for hepatic impairment, neurotoxicity, and cytopenias



Pembrolizumab if MSI high or TMB ≥10

PV Pathogenic Variant





Breast Cancer – Stage IV Any ER/PR and HER2+

If patient ER/PR+ add anastrozole and OS or ADT when only receiving HER2 directed therapy Neratinib and Yes capecitabine Trastuzumab and Perform germline testing vinorelbine Tucatinib, trastuzumab, Ado-trastuzumab and CGP testing Yes and capecitabine d emtansine d Lapatinib and Fam-trastuzumab Candidate capecitabine i Brain metastases? Yes deruxtecan de for chemotherapy? Received Tucatinib. trastuzumab. Trastuzumab and Ado-trastuzumab No adjuvant taxane (Stage IV Any ER/PR and HER2+ abc) emtansine d and capecitabine d chemotherapy fg HER2 directed the rapy within <12 mo? Maintenance trastuzumab Trastuzumab and Trastuzumab, pertuzumab, and No No lapatinib f docetaxel (4-6 mo. max response) d and pertuzumab d Abemaciclib and trastuzumab Clinical trial(s) always considered on pathway. plus fulvestrant if HR+

a If Bone Metastases prescribe zoledronic acid; prescribe denosumabif contraindication; vitamin D and dental evaluation; if symptomatic, refer to Radiation Oncology

If Brain Metastases referral to Radiation Oncology; fam-trastuzumab deruxtecan and tucatinib trastuzumab capecitabine are preferred in patients brain metastases

Fif MSI High or TMB ≥10 pembrolizumab is recommended to be given after endocrine therapy

Evaluate Cardiovascular Risk Factors with baseline LVEF (with ECHO or MUGA) and CMP; monitor LVEF every 3 months during therapy

Fam-trastuzumab-Deruxtecan avoid in pneumonitis, Interstitial Lung Disease (ILD)

Multiple Combinations of HER2 Directed Therapies and chemotherapy are FDA approved but optimal sequencing unknown; consider performance status and toxicity profile

⁹ Chemotherapy includes vinorelbine, docetaxel, carboplatin, eribulin, gemcitabine, capecitabine

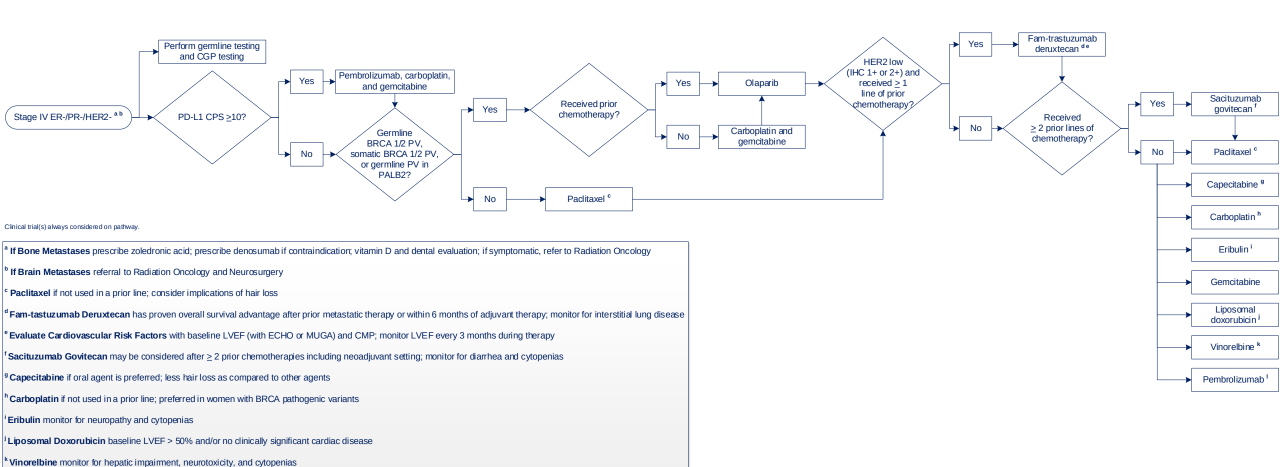
PV Pathogenic Variant







Breast Cancer – Stage IV ER-/PR-/HER2-



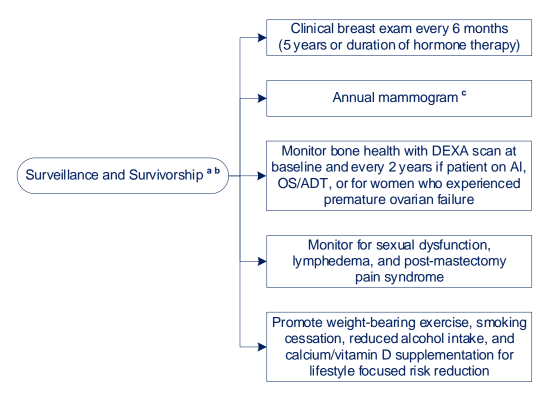


^IPembrolizumab if MSI high or TMB ≥10





Breast Cancer – Surveillance and Survivorship



Clinical trial(s) always considered on pathway.

- ^a Surveillance labs, tumor marker, and systemic imaging not recommended for routine surveillance
- Imaging Following Mastectomy routine imaging of that breast is no longer recommended
- ^c Mammogram routine mammograms are not recommended for men







Breast Cancer – Pathology

Pathology

All results reported in accordance with the CAP Breast Biomarker Reporting Protocol

Tissue Handling Requirements:

Specimen handling slice at 5-10 mm intervals prior to fixation

Cold ischemia time (tissue removal to initiation of fixation) <1 hour

Fixation time 6-72 hours in 10% neutral buffered formalin

Unstained slides used within 6 weeks for ER/PR/HER2 testing

<u>Frozen Sections</u> for sentinel lymph nodes, each gross slice should be no thicker than 2 mm and slices should be embedded in a consistent orientation such that consecutive sections represent tissue separated by no more than 2 mm in the direction of the long axis of the lymph node

Recommended Testing:

DCIS – ER testing only (IHC). Other biomarkers not recommended.

Primary invasive - ER (IHC), PR (IHC), and HER2 (IHC with reflex to FISH for equivocal IHC)

Recurrent/Metastatic – ER (IHC), PR (IHC), and HER2 (IHC with reflex to FISH for equivocal IHC)

Multiple invasive foci – test the largest and highest grade focus of each histologic type

HER2 Interpretation and Reflex:

Negative IHC (0 or 1+) - do NOT reflex

0 – no staining or membrane staining that is incomplete and is faint/barely perceptible and in ≤10% of tumor cells

1+ – incomplete membrane staining that is faint/barely perceptible and in >10% of tumor cells

Equivocal IHC (2+) - REFLEX to FISH

2+ — weak to moderate complete membrane staining in >10% of tumor cells or complete membrane staining that is intense but in \leq 10% of tumor cells Positive IHC (3+) — do **NOT** reflex

3+ - complete membrane staining that is intense and > 10% of tumor cells

HER2 FISH – use dual probe strategy; reflex only if IHC is 2+/equivocal

Negative – an average < 4.0 HER2 signals/cell

Positive - ≥ 6.0 HER2 signals/cell, OR

- ≥ 4.0 HER2 signals/cell AND HER2/CEP17 ratio ≥ 2.0





Breast Cancer – Calculation for the CPS and EG Staging System

Calculation for the CPS & EG Staging System				
Stage/Feature		Points		
Clinical Stage (AJCC staging [1])	0	0		
	IIA	0		
	IIB	1		
	IIIA	1		
	IIIB	2		
	IIIC	2		
Pathologic Stage (AJCC staging [1])	0	0		
	I	0		
	IIA	1		
	IIB	1		
	IIIA	1		
	IIIB	1		
	IIIC	2		
Receptor Status	ER negative [2]	1		
Nuclear Grade [3]	Nuclear grade 3	1		

Used to estimate disease specific survival in patients with breast cancer treated with neoadjuvant chemotherapy. To calculate a score: Add the points for clinical stage, pathologic stage, ER status and nuclear grade to derive a sum between 0 and 6.







Breast Cancer – Molecular Testing Table

Eligibility	Test Category	Test Type	Recommended Vendors	NPOP Coverage	Specimen Type		
All Breast Any Stage	IHC	ER, PR, HER2 (If 2+ reflect to FISH)	GLA/Others	No	Tumor Tissue		
	FISH	HER2 FISH (if HER2 IHC is 2+)	GLA/Others	No	Blood, Tumor Tissue		
	Germline NGS*	Germline breast cancer panel or common hereditary panel (**POC) or referral to CCGS	Fulgent Prevention Genetics	Yes Yes	Saliva, Blood		
Stage Fill, ER+ of PR+/HER2-	IHC	ER, PR, HER2 (If 2+ reflect to FISH)	GLA/Others	No	Tumor Tissue		
	FISH	HER2 FISH (if HER2 IHC is 2+)	GLA/Others	No	Blood, Tumor Tissue		
	Gene Expression/Risk Score Test (21 gene RT-PCR Assay)	21 gene RT-PCR Assay (Oncotype DX 21-gene reoccurrence score) (MammaPrint)	Exact Sciences Biotheranostics	No No	Blood		
	Germline NGS*	Germline breast cancer panel or common hereditary panel (**POC) or referral to CCGS	Fulgent Prevention Genetics	Yes Yes	Saliva, Blood		
	IHC	ER, PR, HER2 (If 2+ reflect to FISH) MMR	GLA/Others Tempus (MMR)	No Yes (MMR when ordered with CGP)	Tumor Tissue		
	FISH	HER2 FISH (if HER2 IHC is 2+)	GLA/Others	No	Blood, Tumor Tissue		
All Metastatic	Somatic NGS	CGP (Solid) CGP Liquid if tissue insufficient/NA	Tempus Foundation Medicine	Yes Yes	Tumor Tissue, Blood		
	Germline NGS*	Germline breast cancer panel or common hereditary panel (**POC) or referral to CCGS	Fulgent Prevention Genetics	Yes Yes	Saliva, Blood		
Stage IV ER+ or PR+, HER2-, Failed Endocrine Therapy, Evaluation for Elacestrant Therapy	Molecular Testing	ESR1 mutation testing	Single gene Mayo (tissue based)	No	Tumor Tissue		
	IHC	ER, PR, HER2 (If 2+ reflect to FISH) PD-L1, 22C3 Clone with CPS Score (pembrolizumab) PD-L1, SP143 Clone (atezolizumab) MMR	GLA/Others Tempus (PD-L1 & MMR) Foundation Medicine (PD-L1)	No Yes (when ordered with CGP) Yes (when ordered with CGP)	Tumor Tissue		
	FISH	HER2 FISH (if HER2 IHC is 2+)	GLA/Others	No	Blood, Tumor Tissue		
	Somatic NGS	CGP (Solid); CGP Liquid if tissue insufficient/NA	Tempus Foundation Medicine	Yes Yes	Tumor Tissue, Blood		
	Germline NGS*	Germline breast cancer panel or common hereditary panel (**POC) or referral to CCGS	Fulgent Prevention Genetics	Yes Yes	Saliva, Blood		
Ductal Carcinoma In Situ	IHC	ER	GLA/Others	No	Tumor Tissue		
	Germline NGS*	Germline breast cancer panel or common hereditary panel (**POC) or referral to CCGS	Fulgent Prevention Genetics	Yes Yes	Saliva, Blood		
* Germline NGS test should include at minimum ATM, BRCA1/2, CDH1, CHEK2, NBN, NF1, PALB2, PTEN, STK11, TP53							

** For genetic online ordering, refer to CCGS page for further details







Questions?

Contact VHAOncologyPathways@va.gov





